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**SILVER-FREE BLACK-AND-WHITE THERMOGRAPHIC
MATERIALS**

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SILVER-FREE BLACK-AND-WHITE THERMOGRAPHIC MATERIALS

FIELD OF THE INVENTION

This invention relates to silver-free black-and-white thermographic materials (“direct thermal” materials) that can provide images having improved tone from the incorporation of color dye-forming couplers, blocked color developing agents, and certain oxidizing agents. This invention also relates to methods of imaging using these thermographic materials.

10

BACKGROUND OF THE INVENTION

Silver-containing thermographic imaging materials (“direct thermal” materials) are non-photosensitive materials that are used in a recording process wherein images are generated by the direct application of thermal energy. These materials have been known in the art for many years and generally 15 comprise a support having disposed thereon one or more imaging layers comprising (a) a relatively or completely non-photosensitive source of reducible silver ions, (b) a reducing composition (usually including a developer) for the reducible silver ions, and (c) a suitable hydrophilic or hydrophobic binder. Thermographic materials are sometimes called “direct thermal” materials in the 20 art because they are directly imaged by a source of thermal energy without any transfer of the energy or image from another material.

In a typical thermographic construction, the image-forming layers are based on silver salts of long chain fatty acids. The preferred non-photosensitive reducible silver source is a silver salt of a long chain aliphatic 25 carboxylic acid having from 10 to 30 carbon atoms, such as behenic acid or mixtures of acids of similar molecular weight. At elevated temperatures, the silver of the silver carboxylate is reduced by a reducing agent whereby a black-and-white image of elemental silver is formed.

30 **Problem to be Solved**

Thermographic materials are imaged by contacting them with the thermal head of a thermographic recording apparatus such as a thermal printer or

thermal facsimile to form a visible image (usually a black-and-white image). Heat generated in the thermal print head can range from 100 to many hundreds of °C. Because the contact between the thermal print head and a given area of the thermographic material is very short (a few milliseconds), the thermographic material never reaches the same temperature as the thermal print head.

5 It is difficult to generate a “neutral” black-and-white silver image in such materials due to the strong dependence of image tone on silver particle size and shape. Typically, the silver image tends to have a yellowish tint. Thus, a fine balancing of toning agents (“toners”) and other components (such as reducing 10 agents and development accelerators) is necessary to provide a desired “neutral” image tone but even then the image tone can change depending upon imaging conditions (that is, temperature and time). The use of toning agents to adjust image tone in thermally developable materials is a common practice as described in early literature such as U.S. Patents 3,080,254 (Grant, Jr.), 3,847,612 15 (Winslow), and 4,123,282 (Winslow), and in more recent publications of which there are hundreds with U.S. Patents 5,599,647 (Defieuw et al.) and 6,146,822 (Asanuma et al.) and EP 1,270,255 (Dooms et al.) being representative.

15 There is a need for better and more predictable control of image tone in thermographic materials that can be imaged under a variety of conditions 20 without the use of silver imaging components.

SUMMARY OF THE INVENTION

This invention provides a silver-free, black-and-white thermographic material comprising a support having thereon at least one imaging 25 layer comprising predominantly a hydrophilic or water-dispersible polymeric latex binder, and further comprising:

- a) a color developing agent precursor that releases a color developing agent when heated to a temperature of at least 80°C, and
- b) a cyan dye-forming color coupler that is capable of reacting 30 with the released color developing agent to produce a cyan dye,
- c) a magenta dye-forming color coupler that is capable of reacting with the released color developing agent to produce a magenta dye,

- d) a yellow dye-forming color coupler that is capable of reacting with the released color developing agent to produce a yellow dye, and
- e) an oxidizing agent that is a hindered-amine N-oxyl that is capable of oxidizing the released color developing agent,

5 the material being substantially free of silver metal or reducible silver ions.

In preferred embodiments, this invention provides a silver-free, black-and-white, non-photosensitive thermographic material that comprises a transparent polymer support having on only one side thereof one or more thermally sensitive imaging layers and an outermost non-thermally sensitive protective layer over the one or more thermally sensitive imaging layers, the one or more thermally sensitive imaging layers comprising predominantly one or more hydrophilic binders, and in reactive association, imaging chemistry consisting essentially of:

- 15 a) a color developing agent precursor that releases a *p*-phenylenediamine color developing agent when heated to a temperature of at least 80°C, the color developing agent precursor being present in an amount of from about 0.001 to about 0.05 mol/m²,
- b) a cyan dye-forming color coupler that is capable of reacting with the released color developing agent to produce a cyan dye,
- 20 c) a magenta dye-forming color coupler that is capable of reacting with the released color developing agent to produce a magenta dye,
- d) a yellow dye-forming color coupler that is capable of reacting with the released color developing agent to produce a yellow dye,
- 25 e) an oxidizing agent that is 2,2,6,6-tetramethylpiperidinyloxy, free radical, 4-hydroxy-2,2,6,6-tetramethylpiperidinyloxy, free radical, or 4-hydroxy-2,2,6,6-tetramethylpiperidinyloxy benzoate, free radical and is present in an amount of from about 1 to about 10 mol/mol of the color developing agent precursor, and
- 30 f) a development enhancing toning agent,

the material being substantially free of silver metal or reducible silver ions, and the cyan dye-forming color coupler, magenta dye-forming color

coupler, and yellow dye-forming color coupler being independently present in an amount from about 0.05 to about 0.5 mol/mol of the color developing agent precursor.

This invention also provides a method comprising imaging the 5 thermographic material of the present invention with a thermal imaging source to provide a visible image.

This method can further include using the imaged thermographic material for medical diagnostic purposes.

When direct thermographic materials are imaged using thermal 10 energy, the conventional components of reducing agent, non-photosensitive silver salt, and toning agents react to form a silver image that may not have the desired color tint or hue (or image tone). However, in the materials of this invention, the blocked color developing agent precursor and dye-forming color couplers provide a combination of cyan, yellow, magenta dyes in appropriate amounts so as to 15 modify the tone of the resulting image. No silver (or silver ions) and conventional black-and-white developer or reducing agents are present. The released color developing agent is oxidized by the unique hindered-amine N-oxyl oxidizing agent. The oxidized color developing agent then reacts with the color couplers to provide appropriate cyan, yellow, and magenta dyes. The resulting image is more 20 nearly neutral in overall density, meaning that the overall red, blue, and green densities are close to each other to the blue density that inherently results from the silver metal. The overall density may be designed to be slightly “blue” in color (i.e., a lower blue density relative to the red and green densities) since users may prefer a bluish-black background for viewing the images.

25 Thus, the present invention provides a more convenient means for adjusting or controlling image tone without the need to use silver imaging components or conventional toning agents of the type that significantly modify silver image tone.

The thermographic materials of this invention comprise a 30 transparent support having thereon an aqueous-based imaging layer(s) comprising predominantly hydrophilic binders such as gelatin or a gelatin derivative, and optionally an aqueous-based or solvent-based overcoat serving as a surface

protective or "slip" layer. Thus, the embodiments of this invention are coated out of aqueous-based formulations.

DETAILED DESCRIPTION OF THE INVENTION

5 The direct thermographic materials of this invention can be used to provide black-and-white images using dye-forming color couplers, blocked color developing agents, specific oxidizing agents, hydrophilic binders, and other components known to be useful in such materials. No silver or silver ions (such as from organic silver salts) are purposely added to the materials.

10 The direct thermographic materials of this invention can be used in black-and-white thermography and in electronically generated black-and-white hardcopy recording. They can be used as output media, in radiographic imaging (for example digital medical imaging), X-ray radiography, and in industrial radiography. Furthermore, the absorbance of these thermographic materials 15 between 350 and 450 nm is desirably low (less than 0.5), to permit their use in the graphic arts area (for example, in image-setting and phototypesetting operations), in the manufacture of printing plates, in contact printing, in duplicating ("duping"), and in proofing.

20 The direct thermographic materials of this invention are particularly useful as output media for medical imaging of human or animal subjects in response to thermal imaging means. Such applications include, but are not limited to, thoracic imaging, mammography, dental imaging, orthopedic imaging, general medical radiography, therapeutic radiography, veterinary radiography, and auto-radiography.

25 In the direct thermographic materials of this invention, the components needed for imaging can be in one or more thermally sensitive layers on one side ("frontside") of the support. The layer(s) that contain the color developing agent precursor, oxidizing agents, and color couplers are referred to herein as thermographic emulsion layer(s) or thermally sensitive imaging layer(s).

30 Where the materials contain thermographic imaging layers on one side of the support only, various non-imaging layers can be disposed on the

“backside” (non-emulsion or non-imaging side) of the materials including an outermost slip layer and/or a conductive layer.

In such embodiments, various non-imaging layers can also be disposed on the “frontside,” imaging, or emulsion side of the support, including 5 primer layers, interlayers, opacifying layers, subbing layers, carrier layers, antihalation layers, “slip” (or protective) layers, auxiliary layers, and other layers readily apparent to one skilled in the art.

For some embodiments, the direct thermographic materials may be “double-sided” or “duplitized” and have thermographic emulsion coating(s) or 10 thermally sensitive imaging layer(s) on both sides of the support. In such constructions each side can also include one or more primer layers, interlayers, antistatic layers, auxiliary layers, conductive layers, “slip” (or protective) layers, and other layers readily apparent to one skilled in the art.

15 Definitions

As used herein:

In the descriptions of the thermographic materials of the present invention, “a” or “an” component refers to “at least one” of that component (for example, a color developing agent precursor or color coupler).

20 “Thermographic material(s)” means a construction comprising at least one thermographic emulsion layer or thermally sensitive imaging layer(s) wherein the required components or optional additives are distributed, as desired, in the same layer or in adjacent coated layers, as well as any supports, topcoat layers, image-receiving layers, carrier layers, blocking layers, conductive layers, 25 antihalation layers, subbing or priming layers. These materials include multilayer constructions in which one or more imaging components are in different layers, but are in “reactive association”. Thus, one layer can include the color developing agent precursor and another layer can include the oxidizing agent, but the two reactive components are in reactive association with each other.

30 When used in thermography, the term, “imagewise exposing” or “imagewise exposure” means that the material is imaged using any means that provides an image using heat. This includes, for example, analog exposure where

an image is formed by differential contact heating through a mask using a thermal blanket or infrared heat source, as well as by digital exposure where the image is formed one pixel at a time such as by modulation of thermal print-heads or laser imaging sources.

5 The materials of this invention are “direct” thermographic materials used in “direct thermal transfer” in which imaging is either “on” or “off” (bimodal), and thermal imaging is carried out in a single “element” containing all of the necessary imaging chemistry. Direct thermal imaging is distinguishable from what is known in the art as thermal transfer imaging (such as dye transfer 10 imaging) in which the image is produced in one element (“donor”) and transferred to another element (“receiver”) using thermal means.

“Catalytic proximity” or “reactive association” means that the components are in the same layer or in adjacent layers so that they readily come into contact with each other during thermal imaging and development.

15 “Emulsion layer,” “imaging layer,” or “thermographic emulsion layer,” means a thermally sensitive layer of a thermographic material that contains the color developing agent precursor. It can also mean a layer of the thermographic material that contains, in addition to this component, additional required components or optional additives. These layers are usually on what is 20 known as the “frontside” of the support.

The slip layer is generally the outermost layer on the imaging side of the material that is in direct contact with the imaging means.

25 Many of the chemical components used herein are provided as a solution. The term “active ingredient” means the amount or the percentage of the desired material contained in a sample. All amounts listed herein are the amount of active ingredient added unless otherwise specified.

30 “Ultraviolet region of the spectrum” refers to that region of the spectrum less than or equal to 410 nm, and preferably from about 100 nm to about 410 nm. “Visible region of the spectrum” refers to that region of the spectrum of from about 400 nm to about 700 nm. “Infrared region of the spectrum” refers to that region of the spectrum of from about 700 nm to about 1400 nm.

“Non-photosensitive” means not intentionally light sensitive. The direct thermographic materials of the present invention are non-photosensitive meaning that no photosensitive silver halide(s) has been purposely added.

The sensitometric terms, absorbance, contrast, D_{min} , and D_{max} have 5 conventional definitions known in the imaging arts. In thermographic materials, D_{min} is considered herein as image density in the non-thermally imaged areas of the thermographic material. The sensitometric term absorbance is another term for optical density (OD).

“Transparent” means capable of transmitting visible light or 10 imaging radiation without appreciable scattering or absorption.

The terms “double-sided”, “double-faced coating”, or “dupliclized” are used to define thermographic materials having one or more of the same or different imaging layers disposed on both sides (front and back) of the support.

As a means of simplifying the discussion and recitation of certain 15 substituent groups, the term “group” refers to chemical species that may be substituted as well as those that are not so substituted. Thus, the term “alkyl group” is intended to include not only pure hydrocarbon alkyl chains, such as methyl, ethyl, *n*-propyl, *t*-butyl, cyclohexyl, *iso*-octyl, and octadecyl, but also alkyl chains bearing substituents known in the art, such as hydroxyl, alkoxy, 20 phenyl, halogen atoms (F, Cl, Br, and I), cyano, nitro, amino, and carboxy. Also, an alkyl group can include ether and thioether groups (for example CH₃-CH₂-CH₂-O-CH₂- and CH₃-CH₂-CH₂-S-CH₂-), haloalkyl, nitroalkyl, alkylcarboxy, carboxyalkyl, carboxamido, hydroxyalkyl, sulfoalkyl, and other groups readily apparent to one skilled in the art.

25 *Research Disclosure* is a publication of Kenneth Mason Publications Ltd., Dudley House, 12 North Street, Emsworth, Hampshire PO10 7DQ England. It is also available from Emsworth Design Inc., 147 West 24th Street, New York, N.Y. 10011.

30 Other aspects, advantages, and benefits of the present invention are apparent from the detailed description, examples, and claims provided in this application.

Color Developing Agent Precursors and Dye-Forming Color Couplers

The present invention uses one or more color developing agent precursors in the thermographic materials. By "precursor" is meant that the compounds are capable of releasing a compound that is a color developing agent when heated to a temperature of at least 80°C. Such precursor compounds may also be described as "blocked" color developing agents that become "unblocked" or reactive upon heating to the appropriate temperature. The released color developing agents can be any of those known in the art for providing color images in color photographic materials including but not limited to, aminophenols, *p*-phenylenediamines (especially N,N-dialkyl-*p*-phenylenediamines) and others which are well known in the art, such those described in EP 0 434 097A1 (published June 26, 1991) and EP 0 530 921A1 (published March 10, 1993). It may be useful for the released color developing agents to have one or more water-solubilizing groups as are known in the art. Further details of such materials are provided in *Research Disclosure*, publication 38957, pages 592-639 (September 1996). The color developing agent precursors then have an appropriate "blocking" group that prohibits there reaction with a dye-forming color coupler until the color developing agent is released during thermal imaging. Useful blocking groups would be readily apparent to one skilled in the art.

Representative color developing agent precursors are described in several publications including U.S. Patent Publication 2002/0018967 (Irving et al.), incorporated herein by reference for the compounds described in paragraphs 0143 through 0228 including the specific compounds identified as D-1 through D-46. These compounds can be prepared using known procedures and starting materials as described in the art.

Particularly useful color developing agent precursors are identified below for use in the Examples as CDA-1, CDA-2, CDA-3, CDA-4, and CDA-5.

The one or more color developing agent precursors are present in an amount of from about 0.0001 to about 0.1 mol/m² and preferably in an amount of from about 0.001 to about 0.05 mol/m².

The photothermographic materials of this invention also include a combination of one or more magenta dye-forming color couplers, one or more

yellow dye-forming color couplers, and one or more cyan dye-forming color couplers to provide the desired neutral images described herein. Any convenient cyan, yellow, and magenta dye-forming color couplers can be employed as would be determined by a skilled worker in the art through routine experimentation to 5 determine how much of what color couplers would improve the desired neutral image tone. In general, the amount of such dye-forming couplers is from about 0.01 to about 1 mol and preferably from about 0.05 to about 0.5 mol, per mole of color developing agent precursor, independently for the cyan dye-forming color couplers, yellow dye-forming color couplers, and the magenta dye-forming color 10 couplers.

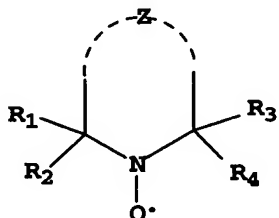
Conventional dye forming couplers are described in considerable publications too numerous to mention including *Research Disclosure*, Number 389, Item 38957, Section X. Dye image formers and modifiers, B. Image-dye-forming couplers, publications noted therein. Representative cyan dye-forming 15 color couplers are described in U.S. Patent 5,453,348 (Kuse et al.). Examples of useful cyan dye-forming color couplers include compounds having a naphthol or phenol structure and that form indoaniline dyes via the coupling reaction with a color developing agent. Representative examples of magenta dye-forming color couplers include compounds having a 5-pyrazolone ring with an active methylene 20 group and pyrazoloazole compounds. Representative examples of yellow dye-forming color couplers have benzoylacetoanilide, pivalylacetoanilide, and acylacetoanilide structures containing active methylene rings. Both 2-equivalent and 4-equivalent dye-forming color couplers can be used. Such color couplers 25 can be prepared using well known procedures and starting materials as described in many publications.

Particularly useful dye-forming color couplers are identified below for the Examples as C-1 (cyan), C-2 (cyan), Y-1 (yellow), and M-1 (magenta).

Oxidizing Agents

30 The thermographic materials also include one or more oxidizing agents that are hindered-amine N-oxyls. These compounds are stable radicals that are capable of oxidizing the released color developing agent but do not

substantially react with other chemical components in the imaging layers. They can be generally defined as having the following Structure I:



5

(I)

wherein R₁, R₂, R₃, and R₄ are independently substituted or unsubstituted alkyl groups, substituted or unsubstituted aryl groups, or substituted or unsubstituted cycloalkyl groups. Such alkyl groups can have 1 to 20 carbon atoms and can be branched or linear (such as methyl, ethyl, *iso*-propyl, *t*-butyl, *n*-hexyl, dodecyl, 10 benzyl, and methoxymethyl groups). Such aryl groups have 6 or 10 carbon atoms in the aromatic ring and can be substituted with one or more alkyl or alkoxy groups (for example, phenyl, naphthyl, 3-methoxyphenyl, and 2,4-dimethylphenyl groups). The cycloalkyl groups have 5 to 10 carbon atoms in the ring structure [such as cyclopentyl, cyclohexyl, 2,4-dimethylcyclohexyl, and 2,4-di(*t*-butyl)cyclohexyl groups]. Preferably, R₁, R₂, R₃, and R₄ are independently substituted or unsubstituted alkyl groups having 1 to 4 carbon atoms, substituted or unsubstituted cyclohexyl groups, or substituted or unsubstituted phenyl groups. More preferably, they are independently unsubstituted methyl, ethyl, cyclohexyl, or phenyl groups. In most preferred embodiments, R₁, R₂, R₃, and R₄ are the same 15 (such as methyl). 20

Alternatively, either R₁ and R₂ or R₃ and R₄ can be combined to form a carbocyclic or heterocyclic ring, including fused ring systems, that can be further substituted with a variety of substituents that do not interfere with the oxidizing capacity of the compounds.

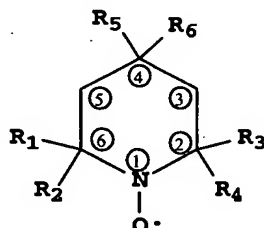
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In Structure I, Z represents the carbon or nitrogen atoms necessary to complete a 5- to 14-membered heterocyclic ring (including fused ring systems). These heterocyclic rings can be further substituted with various groups such as the substituted or unsubstituted alkyl, aryl, and cycloalkyl groups defined above, or

with other groups such as oxo, hydroxy, alkyl esters, aryl esters, and sulfonyl esters.

More specifically, the hindered amine N-oxyls useful in the present invention can be represented by the following Structure II:

5



(II)

wherein R₁, R₂, R₃, and R₄ are defined as described above for Structure I. R₅ and R₆ are independently hydrogen, substituted or substituted alkyl, aryl, or cycloalkyl groups as defined for R₁, R₂, R₃, and R₄. In addition, R₅ and R₆ can be independently hydroxy (or oxo), alkyl esters [that is, -OC(=O)-alkyl groups], aryl esters [that is, -OC(=O)-aryl groups], or sulfonyl esters [that is, -OS(=O)(=O)-alkyl groups]. In addition, R₅ and R₆ can be combined to form an oxo or keto group (=O), or they can be combined to form a substituted or unsubstituted carbocyclic or heterocyclic ring or fused ring system. Preferably, at least one of R₅ and R₆ is hydrogen, and more preferably, at least one is hydrogen and the other is hydroxy, an alkyl group, or an alkyl or aryl ester. Most preferably, one of R₅ and R₆ is hydrogen and the other is hydroxy or benzoate.

While the predominant substituents of the hindered amine N-oxyls are in the 2-, 4-, and 6-positions of Structure II, there may also be additional substituents in the 3- and 5-positions as selected from the various substituents defined above.

Specific examples of useful hindered-amine N-oxyls are 2,2,6,6-tetramethylpiperidinyloxy, free radical (also known as "TEMPO"), 4-hydroxy-2,2,6,6-tetramethylpiperidinyloxy, free radical (also known as "4-hydroxy-TEMPO"), 4-hydroxy-2,2,6,6-tetramethylpiperidinyloxy benzoate, free radical (also known as "4-hydroxy-TEMPO benzoate"), 2,2,6,6,-tetramethyl-4-

(methylsulfonyloxy)-1-piperidinoxy, free radical (also known as “4-hydroxy-TEMPO mesylate”).

The one or more oxidizing agents are present in the thermographic materials in an amount of from about 0.5 to about 20 mol, and preferably from 5 about 1 to about 10 mol, per mol of the color developing agent precursor in the materials.

These oxidizing agents can be obtained from several commercial sources including Aldrich Chemical Company or they can be prepared using known synthetic procedures and starting materials as described for example by 10 Klemchuk, ACS Symposium Series, “Introduction to Hindered Amine Stabilizers”, pp. 1-10, 1985, Toda et al. ACS Symposium Series, “Progress in the Light Stabilization of Polymers”, pp. 37-54, 1985, and by Yamaguchi et al., *Pure & Appl. Chem.* 62(2), pp. 217-222, 1990, Inokuchi et al. *J.Org. Chem.*, 55(2), pp. 462-466, 1990, and Kirchhoff et al., *Polymers & Polymer Composites*, 8(4), pp. 15 245-254, 2000.

Other Addenda

The direct thermographic materials of this invention can also contain other additives such as shelf-life stabilizers, contrast enhancers, dyes or 20 pigments, post-processing stabilizers or stabilizer precursors, thermal solvents (also known as melt formers), and other image-modifying and development-modifying agents as would be readily apparent to one skilled in the art. Toning agents that are commonly used in the art to modify image tone of the reduced silver are not necessary in the practice of this invention because image 25 tone is generated and controlled from the dye-forming color couplers. However, toning agents that accelerate development and/or increase image density may be useful.

Suitable stabilizers that can be used alone or in combination include thiazolium salts as described in U.S. Patents 2,131,038 (Staud) and 30 2,694,716 (Allen), azaindenes as described in U.S. Patent 2,886,437 (Piper), triazaindolizines as described in U.S. Patent 2,444,605 (Heimbach), the urazoles as described in U.S. Patent 3,287,135 (Anderson), sulfocatechols as described in

U.S. Patent 3,235,652 (Kennard), oximes as described in GB 623,448 (Carrol et al.), polyvalent metal salts as described in U.S. Patent 2,839,405 (Jones), thiuronium salts as described in U.S. Patent 3,220,839 (Herz), palladium, platinum, and gold salts as described in U.S. Patents 2,566,263 (Tirelli) and 5 2,597,915 (Damshroder), compounds having $-\text{SO}_2\text{CBr}_3$ groups as described for example in U.S. Patents 5,594,143 (Kirk et al.) and 5,374,514 (Kirk et al.), and 2-(tribromomethylsulfonyl)quinoline compounds as described in U.S. Patent 5,460,938 (Kirk et al.).

10 Stabilizer precursor compounds capable of releasing stabilizers upon application of heat during imaging can also be used. Such precursor compounds are described in for example, U.S. Patent 5,158,866 (Simpson et al.), U.S. Patent 5,175,081 (Krepksi et al.), U.S. Patent 5,298,390 (Sakizadeh et al.), and U.S. Patent 5,300,420 (Kenney et al.).

15 In addition, certain substituted-sulfonyl derivatives of benzotriazoles may be used as stabilizing compounds as described in U.S. Patent 6,171,767 (Kong et al.) and U.S. Patent 6,083,681 (Lynch et al.).

20 The direct thermographic materials of this invention may also include one or more thermal solvents (or melt formers) as disclosed in U.S. Patents 3,438,776 (Yudelson), 5,250,386 (Aono et al.), 5,368,979 (Freedman et al.), 5,716,772 (Taguchi et al.), and 6,013,420 (Windender).

Binders

25 The color developing agent precursor, dye-forming color couplers, hindered-amine N-oxyl oxidizing agents, and any optional additives used in the present invention are generally mixed with one or more hydrophilic binders to form an aqueous-based coating formulation. Water-dispersible polymer latex binders can also be used.

30 Thus, the binders are predominantly (at least 50% by weight of total binders) hydrophilic in nature and aqueous solvent-based formulations are used to prepare such thermographic materials. Mixtures of hydrophilic binders can also be used.

Examples of useful hydrophilic binders that can be used include proteins and protein derivatives, gelatin and gelatin-like derivatives (hardened or unhardened), cellulosic materials, acrylamide/methacrylamide polymers, acrylic/methacrylic polymers polyvinyl pyrrolidones, polyvinyl alcohols, 5 poly(vinyl lactams), polymers of sulfoalkyl acrylate or methacrylates, hydrolyzed polyvinyl acetates, polyacrylamides, polysaccharides, and other synthetic or naturally occurring vehicles commonly known for use in aqueous-based imaging emulsions.

Water-dispersible binders including water-dispersible polymer 10 latexes can also be used in place of some or all of the hydrophilic binders in the thermographic materials of this invention. Such materials are well known in the art including U.S. Patent 6,096,486 (noted above).

In some embodiments, hydrophobic binders can be used as long as they comprise less than 50 weight % of total binder weight. Examples of useful 15 hydrophobic binders include polyvinyl acetals, polyvinyl chloride, polyvinyl acetate, cellulose acetate, cellulose acetate butyrate, polyolefins, polyesters, polystyrenes, polyacrylonitrile, polycarbonates, methacrylate copolymers, maleic anhydride ester copolymers, butadiene-styrene copolymers, and other materials readily apparent to one skilled in the art. Copolymers (including terpolymers) are 20 also included in the definition of polymers. The polyvinyl acetals (such as polyvinyl butyral and polyvinyl formal), cellulose ester polymers, and vinyl copolymers (such as polyvinyl acetate and polyvinyl chloride) are preferred. Particularly suitable binders are polyvinyl butyral resins that are available as 25 BUTVAR® B79 (Solutia, Inc.) and PIOLOFORM® BS-18 or PIOLOFORM® BL-16 (Wacker Chemical Company) and cellulose ester polymers.

The polymer binder(s) is used in an amount sufficient to carry the components dispersed therein. Generally, one or more binders are used at a level of about 10% by weight to about 90% by weight (more preferably at a level of about 20% by weight to about 70% by weight) based on the total dry weight of the 30 layer in which it is included.

Support Materials

The thermographic materials of this invention comprise a polymeric support that is preferably a flexible, transparent film that has any desired thickness and is composed of one or more polymeric materials, depending upon their use. The supports are generally transparent (especially if the material is used as a photomask) or at least translucent, but in some instances, opaque supports may be useful. They are required to exhibit dimensional stability during thermal imaging and development and to have suitable adhesive properties with overlying layers. Useful polymeric materials for making such supports include polyesters, cellulose acetate and other cellulose esters, polyvinyl acetal, polyolefins, polycarbonates, and polystyrenes. Preferred supports are composed of polyesters and polycarbonates.

Support materials can contain various colorants, pigments, and antihalation or acutance dyes if desired. For example, the support can contain conventional blue dyes that differ in absorbance from colorants in the various frontside or backside layers as described in U.S. Patent 6,248,442 (Van Achere et al.). Support materials may be treated using conventional procedures (such as corona discharge) to improve adhesion of overlying layers, or subbing or other adhesion-promoting layers can be used, or treated or annealed to promote dimensional stability.

The thermographic materials preferably have an outermost slip or protective layer on at least the imaging side of the support comprising useful components such as one or more specific lubricants and/or matting agents that are known in the art. The matting agents can be composed of any useful material and may have a size in relation to the slip layer thickness that enables them to protrude through the outer surface of the conductive layer, as described for example, in U.S. Patent 5,536,696 (Horsten et al.). Particularly useful combinations of lubricants are described in copending and commonly assigned U.S.S.N. 10/767,757 (filed on January 28, 2004 by Kenney, Foster, and Johnson) that is incorporated herein by reference.

Thermographic Formulations

An aqueous-based formulation is made in an aqueous solvent that comprises at least 50 volume % water. Some of the components may not be water-soluble and thus may need to be dispersed in organic solvents that are 5 miscible with the solvent used to make the formulation.

The thermographic materials of this invention can be constructed of two or more layers on the imaging side of the support. Two-layer materials would include a single imaging layer and an outermost protective layer. The single imaging layer would contain all of the components needed for imaging, 10 those components desired for the present invention, as well as optional materials such as toning agents, development accelerators, thermal solvents, coating aids, and other additives.

Layers or polymeric materials to promote adhesion in thermographic materials are described for example in U.S. Patents 5,891,610 (Bauer et 15 al.), 5,804,365 (Bauer et al.), 4,741,992 (Przezdziecki), and 5,928,857 (Geisler et al.).

Layers to reduce emissions from the film may also be present as described in U.S. Patents 6,352,819 (Kenney et al.), 6,352,820 (Bauer et al.), and 6,420,102 (Bauer et al.), and in copending and commonly assigned U.S.S.N. 20 10/351,814 (filed January 27, 2003 by Hunt), all incorporated herein by reference.

Layer formulations described herein can be coated by various coating procedures including wire wound rod coating, dip coating, air knife coating, curtain coating, slide coating, or extrusion coating. The formulations can be coated one at a time, or two or more formulations can be coated simultaneously 25 by the procedures described in the art.

When the layers are coated simultaneously using various coating techniques, a "carrier" layer formulation comprising a single-phase mixture of the two or more polymers described above may be used as described in U.S. Patent 6,436,622 (Geisler), incorporated herein by reference.

30 Preferably, two or more layers are applied to a film support using slide coating with the first layer coated on top of the second layer while the second layer is still wet using the same or different solvents (or solvent mixtures).

While the first and second layers can be coated on one side of the film support, manufacturing methods can also include forming one or more layers on the opposing or backside of said polymeric support.

Preferred embodiments include a conductive layer on one or both

5 sides of the support, and more preferably on the backside of the support. Various conductive materials are known in the art such as soluble salts, evaporated metal layers, or ionic polymers as described in U.S. Patent 2,861,056 (Minsk) and U.S. Patent 3,206,312 (Sterman et al.), insoluble inorganic salts as described in U.S. Patent 3,428,451 (Trevoy), electroconductive underlayers as described in U.S.

10 Patent 5,310,640 (Markin et al.), and electrically-conductive metal-containing particles dispersed in a polymeric binder as described in EP 0 678 776A1 (Melpolder et al.). In addition, fluorochemicals such as Fluorad® FC-135 (3M Corporation), ZONYL® FSN (E. I. DuPont de Nemours & Co.), as well as those described in U.S. Patent 5,674,671 (Brandon et al.), U.S. Patent 6,287,754

15 (Melpolder et al.), U.S. Patent 4,975,363 (Cavallo et al.), U.S. Patent 6,171,707 (Gomez et al.), and in copending and commonly assigned U.S. Serial Numbers 10/107,551 (filed March 27, 2002 by Sakizadeh, LaBelle, Orem, and Bhave) and 10/265,058 (filed October 10, 2002 by Sakizadeh, LaBelle, and Bhave) can be used.

20 In preferred embodiments, the conductive layer includes one or more specific non-acicular metal antimonate particles such as non-acicular metal antimonate particles composed of $ZnSb_2O_6$.

Imaging/Development

25 The direct thermographic materials of the present invention can be imaged in any suitable manner consistent with the type of material using any suitable source of thermal energy. The image may be “written” simultaneously with development at a suitable temperature using a thermal stylus, a thermal print head, or a laser, or by heating while in contact with a heat-absorbing material.

30 The thermographic materials may include a dye (such as an IR-absorbing dye) to facilitate direct development by exposure to laser radiation.

Use as a Photomask

The direct thermographic materials of the present invention are sufficiently transmissive in the range of from about 350 to about 450 nm in non-imaged areas to allow their use in a method where there is a subsequent

5 exposure of an ultraviolet or short wavelength visible radiation sensitive imageable medium. The materials may then be used as a mask and positioned between a source of imaging radiation (such as an ultraviolet or short wavelength visible radiation energy source) and an imageable material that is sensitive to such imaging radiation, such as a photopolymer, diazo material, photoresist, or

10 photosensitive printing plate.

In such embodiments, the imaging method of this invention can further comprise:

positioning the imaged thermographic material with the visible image thereon between a source of imaging radiation and an imageable material that is

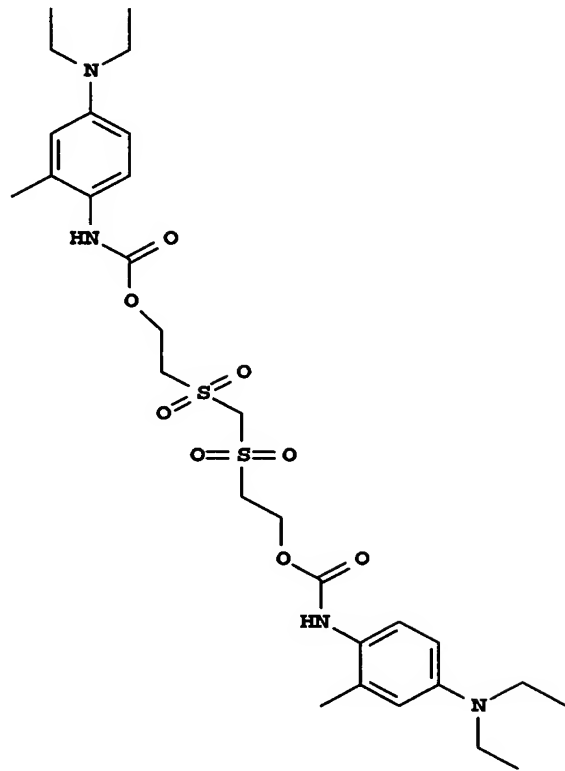
15 sensitive to the imaging radiation, and

thereafter exposing said imageable material to the imaging radiation through the visible image in the imaged thermographic material to provide an image in the imageable material.

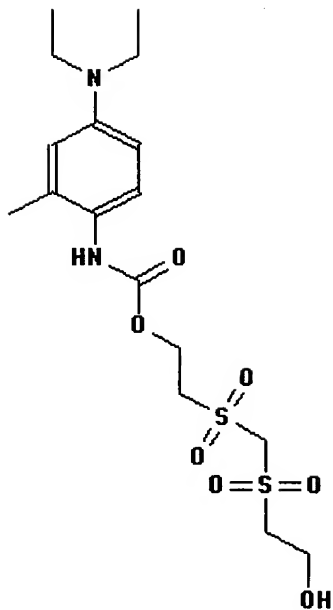
The following examples are provided to illustrate the practice of

20 the present invention and the invention is not meant to be limited thereby.

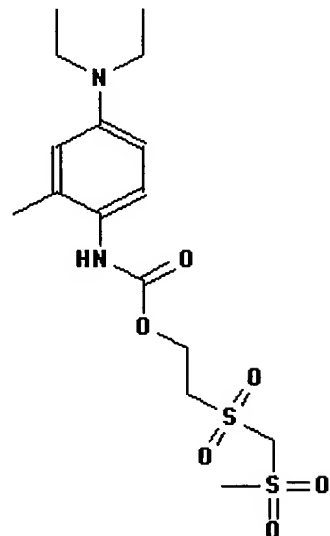
Materials and Methods for the Examples:



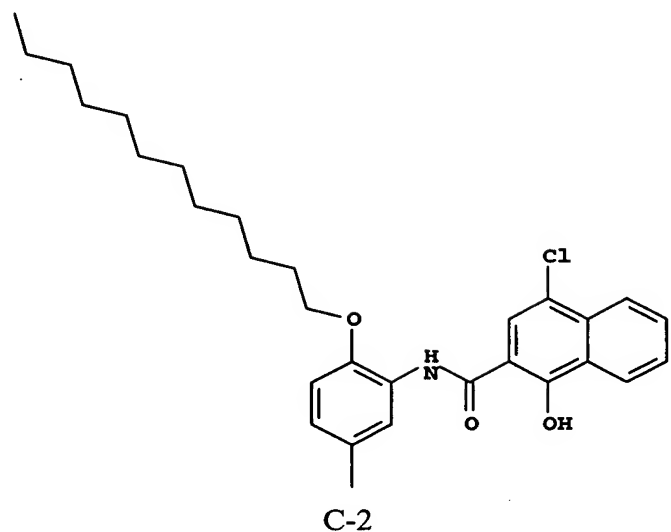
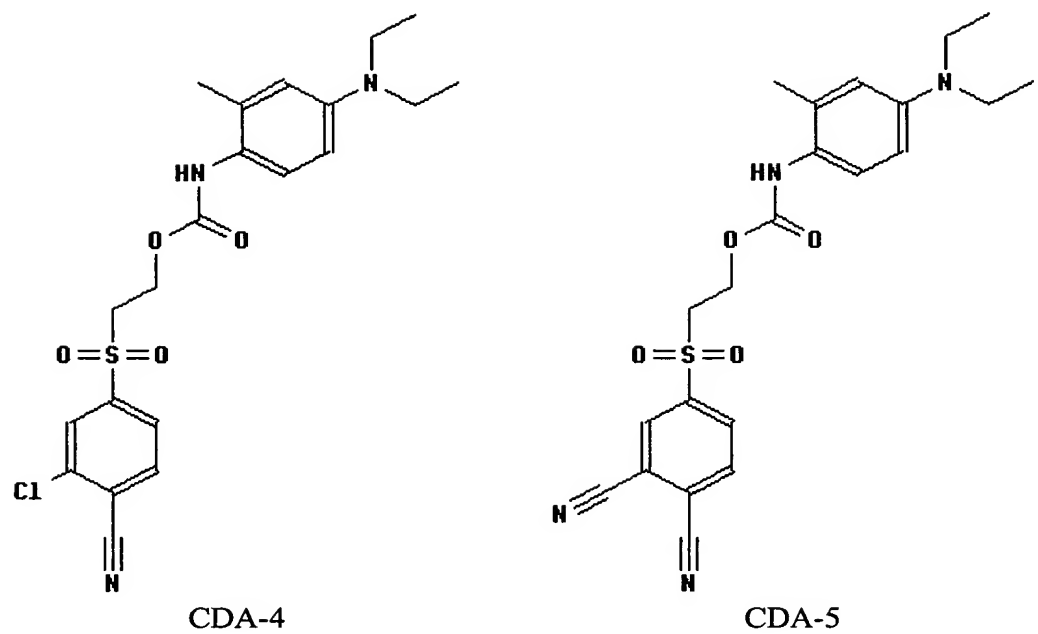
CDA-1

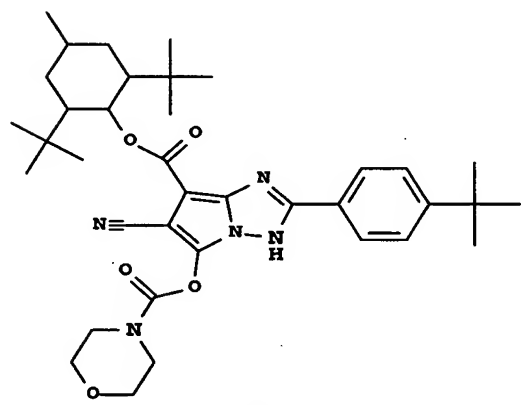
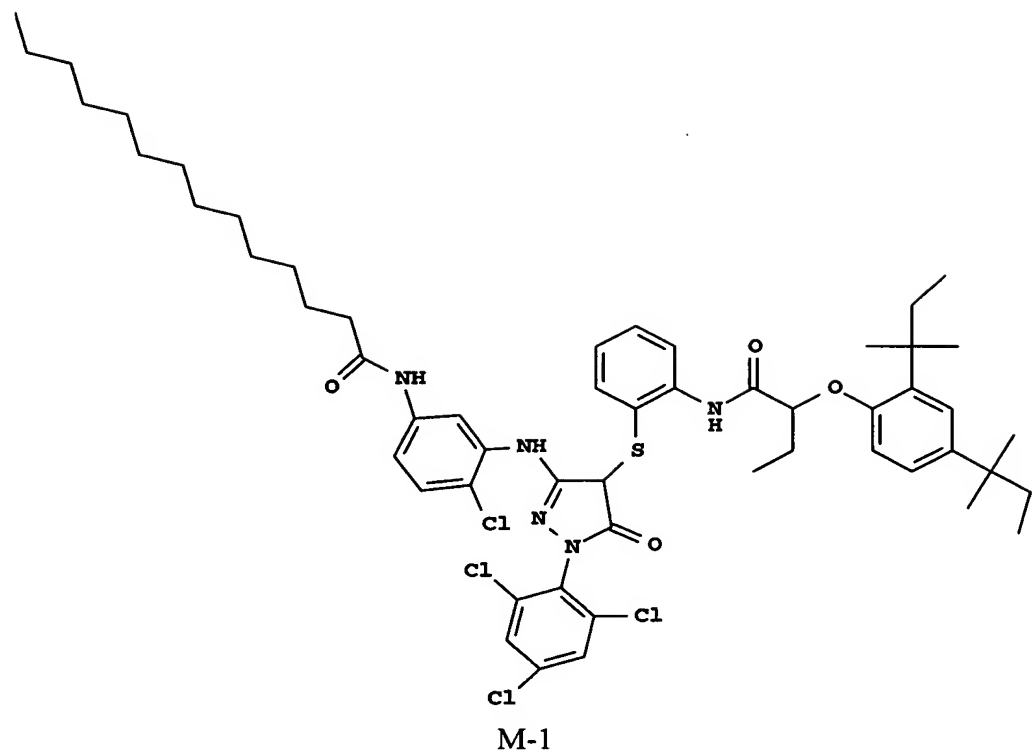


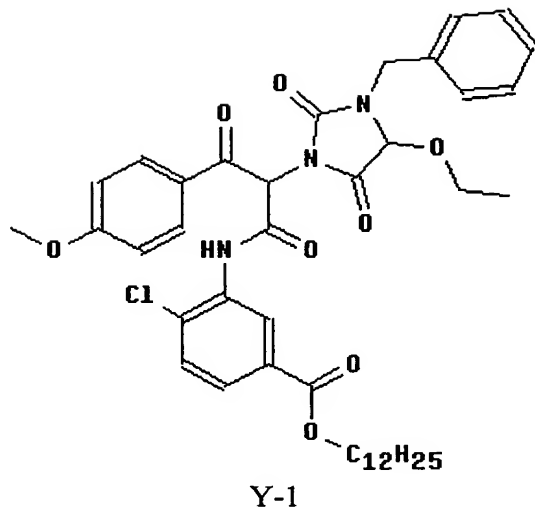
CDA-2



CDA-3







Cyan-1 Coupler Dispersion:

A cyan dye forming coupler dispersion was prepared containing 5 weight % of C-1, 5 weight % of tri(methylphenyl)phosphate (KS1) coupler solvent, and 6 weight % of gelatin using conventional techniques.

Magenta-1 Coupler Dispersion:

A magenta dye forming coupler dispersion was prepared containing 6.8 weight % of M-1, 6.8 weight % of KS1 coupler solvent, and 7.8 weight % of gelatin using conventional techniques.

Yellow-1 Coupler Dispersion:

A yellow dye forming coupler dispersion was prepared containing 9.0 weight % of Y-1, 4.5 weight % of KS1 coupler solvent, and 9 weight % of gelatin using conventional techniques.

Color Developing Agent Precursor Dispersion (Dispersion-1):

A solid particle dispersion of color developing agent precursor was prepared containing 13.2 weight % of CDA-1 and 4 weight % of gelatin.

Color Developing Agent Precursor Dispersion (Dispersion-2):

A solid particle dispersion of color developing agent precursor was prepared containing 10.0 weight % of CDA-4 and 1.0 weight % of Olin 10G surfactant.

HAR1 Hardener Solution:

A hardener composition was prepared containing 2.7 weight % of bis(vinylsulfonyl)methane (BVSM).

SA Dispersion:

5 This was a 25 weight % solid particle dispersion of salicylanilide.

Oxidizing Agent 1:

This compound was 4-hydroxy-2,2,6,6-tetramethylpiperidinyloxy, free radical ("4-hydroxy-TEMPO") that is available commercially from Aldrich Chemical Company.

10 Oxidizing Agent 2:

A solid particle dispersion contained 3.48 weight % of 4-hydroxy-2,2,6,6-tetramethylpiperidinyloxy benzoate, free radical ("4-hydroxy-TEMPO benzoate") that is available commercially from Aldrich Chemical Company.

Oxidizing Agent 3:

15 This compound was 2,2,6,6-tetramethylpiperidinyloxy, free radical ("TEMPO") that is available commercially from Aldrich Chemical Company.

The color densities, both before and after processing are shown in TABLE I provided below. The red, green, and blue densities were measured using Status A densitometry having spectral measuring peaks at 450 nm (for blue density), 550 nm (for green density), and 625 nm (for red density), respectively, using a Macbeth TD504 densitometer and the appropriate filters (see T.H. James, The Theory of the Photographic Process, 4th Ed., Macmillan Publishing Co., Inc., N.Y., 1977, page 521 for details of this process). As the red, green, and blue densities are closer to each other, the more "neutral" are the images. It may be desirable to have a little higher blue density since medical professionals generally prefer to view images in bluish films. Also shown are the average densities of the three color densities and the "spread" that is the maximum difference between the highest and lowest measured color densities. It is desired that the three color densities be high and that the "spread" be low but not necessarily zero depending on the image tone required for a specific imaging material.

Example 1 (Invention):

A direct thermographic material of the present invention was prepared in the following manner:

To 10 g of deionized water, with stirring at 40°C, 1.24 g of 5 oxidized deionized bone gelatin, 1.58 g of Cyan-1 Coupler Dispersion, 1.30 g of Magenta-1 Coupler Dispersion, 1.90 g of Yellow-1 Coupler Dispersion, 5.55 g of Dispersion-1, 1.0 g of SA dispersion, and 0.2 ml of 6.8 weight % SDS solution, were added. The resulting mixture was adjusted to pH 7.0 with a sodium hydroxide solution. Just prior to coating, 0.60 g of Oxidizing Agent 1 and 0.5 ml 10 of HAR1 Hardener solution were added. The resulting formulation was coated at 183 g/m² onto a 0.178 mm gelatin-subbed clear poly(ethylene terephthalate) support. The resulting imaging coating had the following dry component 15 coverage given in g/m²: 14 of gelatin, 0.61 of C-1, 0.68 of M-1, 1.31 of Y-1, 5.62 of CDA-1, 1.92 of salicylanilide, and 4.60 of Oxidizing Agent 1. After drying and hardening the layer for 24 hours, the coated material was cut into 35mm strips (samples) and processed in a thermal processor at 160°C for 18 seconds. The sensitometric results are shown in TABLE I below.

Examples 2-4 (Invention):

20 Thermographic films of this invention were prepared for these examples similarly to that of Invention Example 1 except that the oxidizing agents and amounts were changed as shown in TABLE I below.

Examples 5 (Comparative):

25 Thermographic films outside of the present invention were prepared similarly to that of Invention Example 1 except different oxidizing agents and amounts were used as shown in TABLE I below.

Examples 6-8 (Comparatives):

30 Thermographic films outside of the present invention were prepared similarly to that of Invention Example 1 except that different oxidizing

agents and amounts were used as shown in TABLE I below, and the formulations were coated at 88 g/m².

Examples 9-11 (Comparatives):

5 Thermographic films outside of the present invention were prepared similarly to that of Invention Example 1 except that different oxidizing agents and amounts were used as shown in TABLE I below.

Example 12 (Invention):

10 A thermographic film of the present invention was prepared similarly to that of Invention Example 1 except that a dispersion of Oxidizing Agent 2 was used in place of Oxidizing Agent 1 and with an appropriate amount of water as shown in TABLE I below.

15 **Example 13 (Invention):**

A thermographic film of the present invention was prepared similarly to that of Invention Example 2 except that 1 g of water was used in place of the SA dispersion.

20 **Example 14 (Comparative):**

A thermographic film outside of the present invention was prepared similarly to that of Invention Example 1 except that water was used in place of the oxidizing agent.

25 **Example 15 (Invention):**

To 7.5 g of deionized water, with stirring at 40°C, 1.24 g of oxidized deionized bone gelatin, 1.58 g of Cyan-1 Coupler Dispersion, 1.30 g of Magenta-1 Coupler Dispersion, 1.90 g of Yellow-1 Coupler Dispersion, 9.0 g of Dispersion-2, and 0.2 ml of 6.8 weight % SDS solution, were added. The 30 resulting mixture was adjusted to pH 7.0 with a sodium hydroxide solution. Just prior to coating, 0.30 g of Oxidizing Agent 1 and 0.5 ml of HAR1 Hardener solution were added. The resulting formulation was coated at 183 g/m² onto a

0.178 mm gelatin-subbed clear poly(ethylene terephthalate) support. The resulting imaging coating had the following dry component coverage given in g/m²: 12.5 of gelatin, 0.61 of C-1, 0.69 of M-1, 1.23 of Y-1, 7.0 of CDA-4, and 2.33 of Oxidizing Agent 1. After drying and hardening the layer for 24 hours, the
5 coated material was cut into 35mm strips (samples) and processed in a thermal processor at 160°C for 18 seconds. The sensitometric results are shown in TABLE I below.

TABLE I (Samples Processed at 160°C for 18 seconds)

Example	Oxidizing Agent (g/m ²)	SA (g/m ²)	Densities before Processing Red, Green, Blue	Densities after Processing Red, Green, Blue	Average Density after Processing and (spread)
Invention Example 1	Oxidizing Agent 1 4.60	1.92	0.07, 0.16, 0.18	2.90, 2.94, 3.03	2.96 (0.13)
Invention Example 2	Oxidizing Agent 1 2.30	1.92	0.07, 0.16, 0.19	2.53, 2.49, 2.13	2.38 (0.40)
Invention Example 3	Oxidizing Agent 1 1.53	1.92	0.10, 0.28, 0.24	2.36, 2.42, 1.90	2.23 (0.52)
Invention Example 4	Oxidizing Agent 3 4.17	1.92	0.07, 0.21, 0.18	0.77, 0.84, 0.72	0.78 (0.12)
Comparative Example 5	Sodium persulfate 4.65	1.92	0.13, 0.21, 0.31	0.19, 0.38, 0.47	0.35 (0.28)
Comparative Example 6	Barium peroxide 2.27	0.93	0.18, 0.08, 0.13	0.22, 0.21, 0.33	0.25 (0.12)

Example	Oxidizing Agent (g/m ²)	SA (g/m ²)	Densities before Processing Red, Green, Blue	Densities after Processing Red, Green, Blue	Average Density after Processing and (spread)
Comparative Example 7	Cobalt (III) hexamine trichloride, 2.27	0.93	0.07, 0.13, 0.23	0.14, 0.30, 0.27	0.24 (0.16)
Comparative Example 8	Sodium persulfate 2.27	0.93	0.10, 0.18, 0.22	0.10, 0.19, 0.20	0.16 (0.10)
Comparative Example 9	Phthalazine N-oxide 3.90	1.92	0.11, 0.21, 0.25	0.19, 0.27, 0.56	0.34 (0.37)
Comparative Example 10	4-Picoline N-oxide 2.91	1.92	0.05, 0.11, 0.12	0.16, 0.20, 0.30	0.22 (0.14)
Comparative Example 11	2-Picoline N-oxide 2.91	1.92	0.05, 0.11, 0.14	0.14, 0.17, 0.25	0.19 (0.11)
Invention Example 12	Oxidizing Agent 2 2.67	1.92	0.09, 0.17, 0.22	2.62, 2.65, 2.38	2.55 (0.27)
Invention Example 13	Oxidizing Agent 1 2.30	0	0.05, 0.16, 0.16	2.29, 2.37, 2.18	2.28 (0.19)

Example	Oxidizing Agent (g/m ²)	SA (g/m ²)	Densities before Processing Red, Green, Blue	Densities after Processing Red, Green, Blue	Average Density after Processing and (spread)
Comparative Example 14	None	1.92	0.07, 0.10, 0.14	0.21, 0.24, 0.28	0.24 (0.07)
Invention Example 15	Oxidizing Agent 1 2.33	0	0.07, 0.14, 0.19	1.96, 2.25, 1.96	2.06 (0.29)

The results in TABLE I show that the Invention materials exhibited relatively low density prior to thermal imaging and development and significant image density thereafter. However, the Comparative materials showed poor image densities.

5 The red, green, and blue densities given in TABLE I show that the Invention film samples produced dark and near neutral image tones. Because this invention does not rely on silver metal (known to give unpredictable yellow tone shifts) to make up the image density, the image tone is easily and predictably adjusted by changes in the ratio of the cyan, magenta, and yellow dye-forming
10 color couplers once their relative reactivities and dye color densities have been determined through routine test coatings.

Example 16 (Invention):

 A thermographic material of this invention was prepared similarly
15 to that of Invention Examples 2-4 except that the imaging layer formulation was scaled up and coated to provide 8 x 10 inch (20.3 x 25.4 cm) film sheets. A film sheet was processed in a commercially available AGFA DRYSTAR 2000 resistive thermal head imaging processor to provide an acceptable image of the test pattern.

 The invention has been described in detail with particular reference
20 to certain preferred embodiments thereof, but it will be understood that variations and modifications can be effected within the spirit and scope of the invention.